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(54) Title: CATALYST AND ITS USE IN THE SYNTHESIS OF HYDROGEN PEROXIDE

(57) Abstract: The present invention relates to a catalyst consisting of: (a) one or more metals of the platinum group as active components; (b) one or more polyolefins; and (c) a carrier. The invention also relates to a process for the synthesis of hydrogen peroxide (H_2O_2) from hydrogen and oxygen which uses said catalyst and the use of the hydrogen peroxide solution in oxidation processes catalyzed by titanium silicalite. The process operates under high safety conditions with a high productivity and molar selectivity towards the formation of H_2O_2 .

5 CATALYST AND ITS USE IN THE SYNTHESIS OF HYDROGEN PEROXIDE

The present invention relates to a new catalyst, a process for the direct synthesis of hydrogen peroxide (H_2O_2) from hydrogen and oxygen which uses said catalyst and the use of the hydrogen peroxide solution in oxidation processes catalyzed by titanium-silicalite.

Hydrogen peroxide is a commercially important compound which is widely applied as a bleach in the textile and paper industry, as a biocide in the environmental sector and in the chemical industry in oxidation processes.

Examples of these oxidation processes are those using titanium silicalite as catalysts, such as the epoxidation of olefins (EP 100,119), the ammoximation of carbonyl compounds (U.S. 4,794,198), the oxidation of ammonia to hydroxylamine (U.S. 5,320,819) and the hydroxylation of aromatic hydrocarbons (U.S. 4,369,783).

The industrial production of aqueous solutions of $\rm H_2O_2$ by means of a complex two-step process, is known.

In this process, a solution of anthraquinone, such as

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butylanthraquinone or ethylanthraquinone, in an organic medium immiscible with water, is first hydrogenated and then oxidized with air to produce H_2O_2 which is subsequently extracted in aqueous phase.

This process, however, has substantial disadvantages deriving from the necessity of operating with large volumes of reagents, the numerous steps required, the relatively high cost of the intermediates and the formation of byproducts.

In order to overcome these drawbacks, processes have been studied for the direct synthesis of hydrogen peroxide from H_2 and O_2 . These processes are generally carried out by reacting the two gases in a solvent consisting of an aqueous medium or an aqueous-organic medium, in the pressure of a suitable catalytic system.

Among processes of this type, those which operate in an alcohol or alcohol-aqueous medium, for example in methanol or in methanol-water described, for example, in U.S. patent 4,335,092, in patent application WO 98/16463, in European patent application EP 787681 and more particularly in European patent application EP 978316 and in Italian patent applications MI 2000 A001218, MI 2000 A001219 and MI 2000 A001881, appear attractive from an technical and economic point of view.

With the same conditions in fact, higher reaction

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rates and selectivities have been observed with respect to when operating in an aqueous medium.

The high reaction performances lead in turn to:

- i) the possibility of carrying out the process under high safety conditions, well outside the explosivity zone of H_2 - O_2 mixtures, without jeopardizing it from a technical-economic point of view;
- ii) the possibility of using extremely low quantities of promoters (halides and acids) in the reaction medium,

 with beneficial effects on the stability of the catalytic system and formation of hydrogen peroxide solutions which are stable and at an adequate concentration for direct and economically valid use in oxidation processes.
- 15 Finally, the concentration of the solutions of hydrogen peroxide produced is facilitated up to commercially useful values, as the boiling point and evaporation heat of the suitably selected alcohol are lower than those of water.
- 20 These processes are generally carried out in the presence of a catalytic system consisting of a noble metal, particularly metals of the platinum group or their mixtures, in the form of salts or as supported metals.

It has now been found that it is possible to further 25 improve these processes, in terms of selectivity and cost,

using a heterogeneous catalyst consisting of one or more metals of the platinum group, one or more polyolefins and a carrier.

The use of polyolefins also allows the mechanical properties of the catalyst to be improved and facilitates its filtration from the reaction mixture.

An objective of the present invention therefore relates to a heterogeneous catalyst consisting of one or more metals of the platinum group, one or more polyolefins and a carrier.

Another objective of the present invention relates to a process for the production of hydrogen peroxide starting from hydrogen and oxygen, which uses said catalyst.

A further objective of the present invention relates

15 to the use of solutions of hydrogen peroxide obtained as

described above in an oxidation process catalyzed by tita
nium silicalite.

The catalyst which can be used for the purposes of the invention is a heterogeneous catalyst consisting of:

- 20 (a) one or more metals of the platinum group as active components;
 - (b) one or more polyolefins; and
 - (c) a carrier.

Examples of metals of the platinum group are: palla-25 dium, platinum, ruthenium, rhodium and iridium. Preferred

metals are palladium and platinum.

In these catalysts, the palladium is normally present in a quantity ranging from 0.01 to 4% by weight and the platinum in a quantity ranging from 0.001 to 1% by weight, with an atomic ratio between platinum and palladium ranging from 0.1/99.9 to 50/50.

The palladium is preferably present in a quantity ranging from 0.05 to 2% by weight and the platinum in a quantity ranging from 0.005 to 0.5% by weight, with an atomic ratio between platinum and palladium ranging from 1/99 to 30/70.

In addition to palladium and platinum, other metals such as ruthenium, rhodium, iridium and gold can be present as active components or promoters, in a concentration generally not higher than that of the palladium.

The polyolefins which can be used in the process of the present invention have a molecular weight higher than 400 and are selected from:

- homopolymers of ethylene and copolymers of ethylene with 20 alpha olefins;
 - homopolymers of propylene and copolymers of propylene with alpha olefins;
 - homopolymers of butadiene and copolymers with styrene and other olefins;
- 25 homopolymers of isoprene and copolymers with other ole-

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fins;

- ethylene/propylene (EPR) copolymers;
- ethylene/propylene/diolefin (EPDM) terpolymers;
- thermoplastic elastomers deriving from butadiene and/or
 isoprene and styrene block copolymers, hydrogenated and non-hydrogenated.

The preferred polyolefins are amorphous polyolefins as they are more soluble and therefore easier to disperse on the carrier.

particularly preferred, and, in general, commercial copolymers of butadiene-styrene (synthetic rubber; GRS, SBR); ethylene-propylene (EPM, EPR) copolymers, ethylene-propylene copolymers (EPDM rubbers), styrene-butadiene-styrene (SBR thermoplastic rubbers); isobutylene-isoprene (butyl rubbers).

The polyolefins which can be used for the purposes of the present invention can be prepared according to any of the methods known in the art.

20 The quantity of polyolefin used ranges from 0.1 to 20% by weight, preferably from 1 to 10% by weight, with respect to the catalyst.

The inert carrier may typically consist of activated carbon, silica, alumina, silica-alumina, zeolites, and other materials well known in the state of the art. Acti-

vated carbon is preferably used for the preparation of the catalysts useful for the invention.

Activated carbons which can be used for the purposes of the invention are selected from those of a fossil or natural origin deriving for example from wood, lignite, peat or coconut and having a surface area higher than 100 $\rm m^2/g$, preferably higher than 300 $\rm m^2/g$ and a carbon with a surface area higher than 600 $\rm m^2/g$ is particularly preferred. Preferred activated carbons are those with a low ash content.

The sulfonated activated carbons described in European patent application EP 978316 can be used for the purpose.

Before the deposition of the metals or polyolefins, the activated carbon can be subjected to treatment such as washing with distilled water or treatment with acids, bases or diluted oxidizing agents, for example acetic acid, hydrochloric acid, sodium carbonate and hydrogen peroxide.

The catalyst can be prepared by dispersing the active components on the inert carrier or pretreated with the polyolefin, by means of precipitation and/or impregnation starting from precursors consisting for example of solutions of their salts or soluble complexes, and then reduced to the metallic state by means of thermal and/or chemical treatment with reducing substances such as hydrogen, sodium formiate, sodium citrate or by preparative techniques well

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known in the art.

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According to an embodiment of the present invention, the catalyst can be prepared by dispersing in sequence and alternating the precursors of the single metal components of the catalyst onto the carrier, as described and claimed in patent application IT MI2000-A001219.

The polyolefin is generally dissolved in a suitable solvent and the resulting solution is used to impregnate the carrier.

The dry impregnation technique is preferably used, which consists in putting the polyolefin mixture in contact with the carrier, in a closed reactor at 100-120°C, for 2-3 hours, in order to facilitate the depositing of the polymer onto the surface. At the end, the solvent is evaporated at a temperature of 140°C for 3-4 hours.

Examples of solvents suitable for the purposes of the present invention are selected from paraffins, aromatic hydrocarbons and cyclo-paraffins. N-heptane, toluene, decaline, n-decane are preferably used.

In the preparation of the catalyst, the sequence with which the metals of the platinum group (a) and polyolefins (b) are put in contact with the carrier, is not particularly critical.

When a quantity of polyolefins higher than 5% by 25 weight is used, however, it is preferable to disperse the

mixture of polyolefins onto the carrier after depositing the metals forming the active phase.

The catalyst of the present invention is particularly advantageous in a process for the preparation of hydrogen peroxide from hydrogen and oxygen in a reaction solvent containing a halogenated promoter and/or an acid promoter.

The catalyst is normally dispersed in the reaction medium at a concentration ranging from 0.1 to 10% by weight, preferably from 0.3 to 3% by weight with respect to the reaction solvent.

The reaction solvent consists of one or more alcohols or an alcohol-water mixture to which an aliphatic ether and/or one or more C_5-C_{32} hydrocarbons are optionally added.

Examples of alcohols suitable for the purposes of the present invention are selected from those with from 1 to 6, preferably from 1 to 4, carbon atoms.

Among C_1 - C_4 alcohols, methanol, ethanol, terbutanol (TBA) or their mixtures, are preferred. Methanol is particularly preferred. Among the preferred mixture is a mixture of methanol and water.

The quantity of alcohol(s) ranges from 30 to 99% by weight with respect to the mixtures, preferably from 50 to 98% by weight.

The aliphatic ethers are selected from those having 25 general formula (I)

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$R-O-R_1$ (I)

wherein R and R_1 , the same or different, are alkyl groups with from 1 to 6 carbon atoms. In the compounds having formula (I), R is preferably methyl and R_1 a ter-alkyl.

5 Methyl-terbutylether (MTBE) is particularly preferred.

The quantity of ethers which is used in the solvent mixture depends on the type of alcohol(s) used and generally ranges from 0 to 70% by weight, preferably from 10 to 60% by weight, with respect to the reaction solvent.

10 According to an embodiment of the process of the present invention, the reaction solvent can also contain one or more C_5-C_{32} hydrocarbons.

These hydrocarbons are generally selected from paraffins, cyclo-paraffins or aromatic compounds.

Examples of paraffinic hydrocarbons are preferably selected from those having from 5 to 18 carbon atoms, and can be linear or branched.

Examples of said paraffinic hydrocarbons are n-hexane, n-heptane, n-octane, n-decane or their branched isomers.

Examples of cyclo-paraffinic hydrocarbons are cyclo-hexane, decaline or their derivatives substituted with one or more alkyl groups with from 1 to 6 carbon atoms. Typical examples of these compounds are methyl-cyclohexane, ethyl-cyclohexane or dimethyl-cyclohexane.

25 Aromatic hydrocarbons suitable for the purposes of the

present invention are preferably selected from those having from 6 to 24 carbon atoms.

Examples of aromatic hydrocarbons are benzene, naph-thalene, alkylbenzenes and alkylnaphthalenes with one or more alkyl chains, linear or branched, having from 1 to 18, preferably from 6 to 12 carbon atoms.

Examples of alkylbenzenes are toluene, xylenes (ortho, meta and para), ethylbenzene and cumene.

The quantity of hydrocarbons which is used in the reaction depends on the type of alcohol(s) used and generally ranges from 0 to 20% by weight, preferably from 0.1 to 10% by weight, with respect to the reaction solvent.

The acid promoter may be any substance capable of generating H⁺ hydrogen ions in the reaction solvent and is generally selected from inorganic acids such as sulfuric, phosphoric, nitric acid or from organic acids such as sulfonic acids. Sulfuric acid and phosphoric acid are preferred.

The concentration of the acid generally ranges from 20 to 1000 mg per kg of reaction solvent and preferably from 50 to 500 mg per kg of reaction solvent.

The halogenated promoter can be any substance capable of generating halide ions in the reaction solvent. Substances capable of generating bromide ions are preferred. These substances are generally selected from hydrobromic

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acid and its salts soluble in the reaction medium, for example sodium bromide, potassium bromide, ammonium bromide or sodium bromate. Hydrobromic acid, sodium bromide and potassium bromide are particularly preferred.

The concentration of the halogenated promoter generally ranges from 0.1 to 50 mg per kg of reaction solvent and preferably from 1 to 10 mg per kg of reaction solvent.

The production of hydrogen peroxide is carried out by reacting oxygen and hydrogen in the reaction solvent in the presence of the catalyst and promoters and in the presence or absence of an inert gas selected from nitrogen, helium, argon. Nitrogen is the preferred gas.

The molar ratio H_2/O_2 in the feeding ranges from 1/1 to 1/100, preferably from 1/2 to 1/15 and the concentration of hydrogen in the gaseous phase in contact with the reaction solvent is conveniently maintained at a value lower than 4.5% molar, outside the explosivity limits of the mixture consisting of H_2 , O_2 and, optionally, an inert gas.

According to an embodiment of the process of the pres-20 ent invention, the reaction can be carried out using air instead of pure oxygen.

The reaction is typically carried out at temperatures ranging from -5° to 90°C, preferably from 2 to 50°C, a temperature ranging from 20 to 40°C is particularly preferred, and at a total pressure higher than atmospheric pressure,

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preferably ranging from 10 to 300 bars, 30-100 bars being particularly preferred.

The process according to the present invention can be carried out batchwise or, preferably, in continuous using a reactor suitable for the purpose and selected from those described in the state of the art.

Operating under the above conditions, it is possible to produce hydrogen peroxide under safety conditions with a reaction productivity normally ranging from 30 to 200 g of H_2O_2 (expressed as H_2O_2 at 100%) per litre of reaction medium per hour and with a molar selectivity towards the formation of H_2O_2 , referring to the hydrogen used up, ranging from 60% to 90%.

The solutions of hydrogen peroxide thus obtained can be used directly in oxidation processes which comprise the use of H_2O_2 without onerous intermediate processing such as the removal of acids and solvents.

Furthermore, the process of the present invention is suitable for the production of aqueous solutions of $\rm H_2O_2$ having a commercial titer, by the removal of the organic components from the reaction medium, for example by distillation, which can be recycled to the synthesis.

The process of the present invention allows the reagents to be transformed into H_2O_2 with high conversions and selectivities, obtaining H_2O_2 solutions without acidity or

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containing only traces of acidity and/or salts.

The following examples, which have the sole purpose of describing the present invention in greater detail, should in no way be considered as limiting its scope.

5 EXAMPLE 1

Treatment of the activated carbon carrier

150 g of activated carbon of maritime pine charcoal in powder form (CECA/2S/E) and 1500 ml of distilled water are charged into a 2 liter jacketed glass flask, thermostat-regulated with an oil-heated bath and equipped with a cooler and stirring. After 2 hours at 80°C, the activated carbon is filtered and washed with distilled water.

The activated carbon, still damp, is then charged into the 2 liter glass flask described above and after adding 1500 ml of a solution at 5% by weight of HCl, the temperature is brought to 80°C. After about 2 hours, the mixture is cooled and the activated carbon is washed on a filter with distilled $\rm H_2O$ until the chlorides have been eliminated. The washed activated carbon is recovered and dried in an oven at 120°C for 3 hours.

EXAMPLE 2 (comparative)

Preparation of the catalyst Pt-Pd/C (EX2)

(a) 900 ml of distilled water, 2.8 g of Na_2CO_3 and subsequently 80 g of activated carbon prepared in Example 1, are charged into a glass reactor, having the character-

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istics described in Example 1. The suspension is maintained at room temperature (20-25°C), under stirring, for 10 minutes.

A solution of 8 g of Na₂PdCl₄ at 10% by weight of Pd in 100 ml of distilled water are subsequently added drop-wise over a period of about 10 minutes and the resulting suspension is kept at room temperature for 10 minutes and is then heated in a water bath for 10 minutes to 90°C. A solution containing 0.76 g of sodium formiate in 100 ml of distilled water is then added and the stirring is continued at 90°C for 2 hours.

After cooling to room temperature, the suspension is filtered and the recovered catalyst is washed with distilled water until the chlorides have been eliminated and dried in an oven at 120°C for 3 hours.

(b) the catalyst thus obtained is put in the 2 litre reactor described above and treated following the procedure described in (a), but using a solution of 0.404 g of H_2PtCl_6 (8% by weight of Pt) instead of the solution of Na_2PdCl_4 .

After drying at 120°C, a catalyst is obtained (EX2) containing 0.97% of Pd and 0.038% of Pt on activated carbon.

EXAMPLE 3

25 Preparation of activated Carbons functionalized with sul-

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fonic groups (C-SO₃H)

80 g of activated carbon prepared as described in Example 1 are charged into a jacketed 2 litre glass reactor, thermostat-regulated with an oil-heated bath and equipped with a cooler and stirring, and 240 g of $\rm H_2SO_4$ at 96% are added dropwise in 20 minutes. After homogenizing the mixture with slight stirring, it is heated to 140°C for 2 hours.

The mixture is cooled to room temperature and 200 g of crushed ice (from distilled water) are added in 10 minutes, the mixture is left to cool, a further 1000 ml of distilled water are added, the contents of the reactor are recovered and filtered. The activated carbon thus treated is washed until the sulfate ion in solution has been eliminated.

Upon elemental analysis, the activated carbon thus treated contains 0.38% of S.

EXAMPLE 4 (comparative)

Preparation of the catalyst Pd-Pt/C-SO₃H (EX4)

The same procedure is adopted as described in Example 20 2, but using as carrier 8 g of activated carbon functionalized as specified in Example 3. The catalyst (EX4) is obtained, which upon analysis contains: 0.039% Pt; 0.98% Pd; 0.35% S.

EXAMPLE 5

25 Preparation of the catalyst Pt-Pd/C+2.9%polySTY

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8 g of the catalyst EX2 are charged into a 200 ml glass vacuum flask and flushed with nitrogen for 15 minutes to eliminate the air.

0.24 g of polystyrene (average MW 120000) are dissolved at 70°C in a 100 ml Erlenmeyer flask containing 24 g of toluene. The resulting solution is then added dropwise in 5 minutes, maintaining the stream of nitrogen, into the flask containing the catalyst EX2. The flask containing the catalyst to which the polymer has been added is put under a rotating evaporator and is slowly reflux heated to 110°C for 3 hours. The toluene is distilled under a slight vacuum, the catalyst is poured into a 100 ml glass and dried in an oven at 140°C for 3 hours.

The catalyst (EX5) is obtained, containing upon analy15 sis: (Pt-Pd/C)+2.9%polySTY (0.036% Pt; 0.94% Pd).

EXAMPLE 6

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Preparation of the catalyst Pt-Pd/C-SO₃H+3%polySTY (EX6)

The same procedure is adopted as described in Example 5, but using the catalyst EX4 prepared in Example 4.

The catalyst (EX6) is obtained containing upon analysis: $Pt-Pd/C-SO_3H+3%polySTY$ (0.035% Pt; 0.93% Pd and 0.36% S).

EXAMPLE 7

Preparation of the catalyst Pt-Pd/C (C2-co-C3) (EX8)

The same procedure is adopted as described in Example

5, but using 0.24 g of 65% ethylene-35% propylene copolymer (EPR rubber) with an average molecular weight of 110000 instead of the polystyrene. The catalyst is obtained (EX8) containing upon analysis: $Pt-Pd/(C+2.9\%C_2-co-C_3)$ (0.036% Pt; 0.94% Pd).

EXAMPLE 8

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Preparation of the catalyst Pt-Pd/C-SO₃H(C₂-co-C₃) (EX10)

The same procedure is adopted as described in Example 7, but using 8 g of the catalyst EX4 of Example 4.

The catalyst is obtained (EX10) containing upon analysis: $Pt-Pd/(C-SO_3H+2.9\%C_2-co-C_3)$ (0.036% Pt; 0.95% Pd; 0.37% S).

EXAMPLES 9-10

Examples 5 and 6 are repeated using 0.24 g of butadiene-styrene copolymer (SBR: 75/25) with an average molecular weight of 120000 instead of the C_2 - C_3 copolymer.

The catalysts (EX12 and EX14) are obtained respectively, containing upon analysis:

EX 12: Pt-Pd/(C +2.9%SBR) (0.036% Pt; 0.95% Pd).

EX 14: Pt-Pd/(C-SO₃H+2.9%SBR) (0.036% Pt; 0.94% Pd; 0.36% S).

EXAMPLES 11-18

Synthesis of hydrogen peroxide

A micropilot plant is used, consisting of a Hastelloy C autoclave having a volume of 400 ml, equipped with a thermostat-regulation system, a magnetic drag stirring sys-

tem, a regulation and control system of the pressure during the reaction, a filter for continuously removing the liquid phase containing the reaction products, a feeding system of the mixture of solvent and promoters in which the reaction takes place, a feeding system of the gaseous reagents and a series of regulation and control instruments.

The reaction trend is followed by continuously analyzing the hydrogen and oxygen in the feeding and at the outlet of the reactor.

The concentration of H_2O_2 which is formed is determined in the liquid effluent of the reactor by titration with potassium permanganate. The selectivity with respect to the converted hydrogen is calculated on the basis of the concentration of H_2O_2 in the reaction effluent and on the basis of the analysis of the H_2 leaving the reactor, once the stationary state has been reached.

1.0 g of catalyst prepared as described in examples 2, 4 and 5-8 and 100 g of methanol:water solution (97/3 by weight) containing 6 ppm of HBr (6 mg/kg) and 200 ppm of H_2SO_4 (200 mg/kg) are charged into the reactor.

The autoclave is pressurized, without stirring, at 60 bars with a gaseous mixture consisting of 3.6% of H_2 , 11% of O_2 and 85.4% of N_2 . The stirring is then started up to 800 revs/minute, the pressure is maintained with a continuous stream, 916 normal liters (N1), of the same gaseous

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mixture, with the contemporaneous feeding of 400 g/hour of a methanol:water solution having the composition defined above and containing 6 ppm of HBr and 200 ppm of $\rm H_2SO_4$. The temperature inside the reactor is maintained at 25°C.

The results obtained after 50 hours of reaction are indicated in Table 1.

Table 1

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Catalyst	H ₂ O ₂ %	Selectivity %
EX2 Pt-Pd/C	5.3	64
EX4 Pt-Pd/C-SO ₃ H	5.7	70
EX5 Pt-Pd/C + 3% polysty	5.6	73
EX6 Pt-Pd/C-SO ₃ H + 3% polysty	5.9	80
EX8 Pt-Pd/(C+(C ₂ -co-C ₃))	5.6	71
EX10 Pt-Pd/(C-SO ₃ H+(C ₂ -co-C ₃))	6.0	78
EX12 Pt-Pd/(C+SBR(25-75))	5.8	74
EX14 Pt-Pd/((C-SO ₃ H(SBR(25-75))	5.8	81

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CLAIMS

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 A catalyst useful for the synthesis of hydrogen peroxide starting from hydrogen and oxygen consisting of:

- (a) one or more metals of the platinum group as active components;
- (b) one or more polyolefins; and
- (c) a carrier:
- 2. The catalyst according to claim 1, wherein the metal components of the catalyst are selected from palladium, platinum, ruthenium, rhodium and iridium.
 - 3. The catalyst according to claim 2, wherein the metal components of the catalyst are palladium and platinum.
- 4. The catalyst according to claim 3, wherein the quantity of palladium ranges from 0.01 to 4% by weight and the quantity of platinum ranges from 0.001 to 1% by weight, with an atomic ratio platinum/palladium ranging from 0.1/99.9 to 50/50.
- 5. The catalyst according to claim 4, wherein the quantity of palladium ranges from 0.05 to 2% by weight and the quantity of platinum ranges from 0.005 to 0.5% by weight, with an atomic ratio platinum/palladium ranging from 1/99 to 30/70.
- The catalyst according to claim 3, wherein in addition to palladium and platinum other metals such as, for
 example, ruthenium, rhodium, iridium and gold can also

be present as active components or promoters, in a concentration not higher than that of the palladium.

- 7. The catalyst according to claim 1, wherein the polyolefins have a molecular weight higher than 400 and are selected from:
 - homopolymers of ethylene and copolymers of ethylene with alpha olefins;
 - homopolymers of propylene and copolymers of propylene with alpha olefins;
- homopolymers of butadiene and copolymers with styrene and other olefins;
 - homopolymers of isoprene and copolymers with other olefins;
 - ethylene/propylene (EPR) copolymers;
- ethylene/propylene/diolefin (EPDM) terpolymers;
 - thermoplastic elastomers deriving from butadiene and/or isoprene and styrene block copolymers, hydrogenated and non-hydrogenated.
- 8. The catalyst according to claim 7, wherein the polyolefins are amorphous polyolefins.
 - 9. The catalyst according to claim 7, wherein the polyolefins are selected from rubbers and copolymers of butadiene-styrene (synthetic rubber; GRS, SBR); ethylene-propylene (EPM, EPR) copolymers, ethylene-propylene-diene copolymers (EPDM rubbers), styrene-

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butadiene-styrene (SBR thermoplastic rubbers); isobutylene-isoprene (butyl rubbers).

- 10. The catalyst according to claim 1, wherein the quantity of polyolefin ranges from 0.1 to 20% by weight with respect to the catalyst.
- 11. The catalyst according to claim 10, wherein the quantity of polyolefin ranges from 1 to 10% by weight with respect to the catalyst.
- 12. The catalyst according to claim 1, wherein the carrier

 10 is selected from activated carbon, activated carbon
 functionalized with sulfonic groups, silica, alumina,
 silica-alumina and zeolites.
 - 13. The catalyst according to claim 12, wherein the carrier is an activated carbon with a low ash content and a surface area higher than $100 \text{ m}^2/\text{g}$.
 - 14. The catalyst according to claim 13, wherein the activated carbon has a surface area higher than 300 m^2/g .
 - 15. The catalyst according to claim 14, wherein the activated carbon has a surface area higher than $600~\text{m}^2/\text{g}$.
- 20 16. The catalyst according to claim 1, prepared by dispersing the active components on an inert carrier or pretreated with polyolefins by means of precipitation and/or impregnation.
- 17. The catalyst according to claim 16, prepared by dis-25 persing in sequence and alternating the precursors of

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the single metal components of the catalyst on an inert carrier or pretreated with polyolefins.

- 18. A process for the production of hydrogen peroxide from hydrogen and oxygen in a reaction solvent containing a halogenated promoter and/or an acid promoter, in the presence of a catalyst according to claims 1 to 17.
- 19. The process according to claim 18, wherein the reaction solvent consists of one or more alcohols or a mixture of alcohol-water optionally containing an aliphatic ether and/or one or more C₅-C₃₂ hydrocarbons.
- 20. The process according to claim 19, wherein the alcohol is selected from those having from 1 to 6 carbon atoms.
- 21. The process according to claim 20, wherein the alcohol is selected from those having from 1 to 4 carbon atoms.
 - 22. The process according to claim 21, wherein the alcohol is selected from methanol, ethanol, terbutanol (TBA) or their mixtures.
- 20 23. The process according to claim 22, wherein the alcohol is methanol.
 - 24. The process according to claim 19, wherein the quantity of alcohol(s) ranges from 30 to 99% by weight with respect to the mixture.
- 25 25. The process according to claim 24, wherein the quan-

tity of alcohol(s) ranges from 50 to 98% by weight with respect to the mixture.

26. The process according to claim 19, wherein the aliphatic ether is selected from those defined by general formula (I)

$R-O-R_1$

wherein R and R_1 , the same or different, are alkyl groups having from 1 to 6 carbon atoms.

- 27. The process according to claim 26, wherein in the compounds having formula (I), R is methyl and R_1 a teralkyl.
 - 28. The process according to claim 27, wherein the ether is methyl-terbutylether (MTBE).
- 29. The process according to claim 19, wherein the quantity of aliphatic ether having general formula (I) ranges from 0 to 70% by weight with respect to the reaction solvent.
- 30. The process according to claim 29, wherein the quantity of aliphatic ether having general formula (I) ranges from 10 to 60% by weight with respect to the reaction solvent.
 - 31. The process according to claim 19, wherein the C_5-C_{32} hydrocarbons are selected from paraffins, cycloparaffins and aromatic compounds.
- 25 32. The process according to claim 31, wherein the paraf-

finic hydrocarbons can be linear or branched.

33. The process according to claim 32, wherein the paraffinic hydrocarbons are selected from those having from 5 to 18 carbon atoms.

- 5 34. The process according to claim 33, wherein the paraffinic hydrocarbons are selected from n-hexane, nheptane, n-octane, n-decane or their branched isomers.
 - 35. The process according to claim 31, wherein the cycloparaffinic hydrocrabons are selected from cyclohexane, decaline or their derivatives substituted with one or more alkyl groups having from 1 to 6 carbon atoms.
 - 36. The process according to claim 35, wherein the substituted cyclo-paraffins are selected from methyl-cyclohexane, ethyl-cyclohexane and dimethyl-cyclohexane.
 - 37. The process according to claim 31, wherein the aromatic hydrocarbons are selected from those having from 6 to 24 carbon atoms.
- 38. The process according to claim 37, wherein the aromatic hydrocarbons are selected from benzene, naphthalene, alkylbenzenes and alkylnaphthalenes with one or more linear or branched alkyl chains having from 1 to 18 carbon atoms.
- 39. The process according to claim 38, wherein the alkyl-25 benzenes and alkylnaphthalenes have a linear or

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branched alkyl chain having from 6 to 12 carbon atoms.

40. The process according to claim 39, wherein the alkylbenzenes are selected from toluene, xylenes (ortho, meta and para), ethylbenzene and cumene.

- 5 41. The process according to claim 19, wherein the quantity of hydrocarbons ranges from 0 to 20% by weight with respect to the reaction solvent.
 - 42. The process according to claim 41, wherein the quantity of hydrocarbons ranges from 0.1 to 10% by weight with respect to the reaction solvent.
 - 10% by weight with respect to the reaction solvent.
- 44. The process according to claim 43, wherein the cata
 lyst is used at a concentration ranging from 0.3 to 3%

 by weight with respect to the reaction solvent.
 - 45. The process according to claim 18, wherein the acid promoter is selected from substances capable of generating H⁺ hydrogen ions in the reaction solvent.
- 20 46. The process according to claim 45, wherein the acid promoter is selected from inorganic acids such as sulfuric, phosphoric, nitric acid or from organic acids such as sulfonic acids.
- 47. The process according to claim 46, wherein the acid promoter is sulfuric acid or phosphoric acid.

48. The process according to claim 18, wherein the concentration of acid promoter ranges from 20 to 1000 mg per kg of reaction solvent.

- 49. The process according to claim 48, wherein the concentration of acid promoter ranges from 50 to 500 mg per kg of reaction solvent.
 - 50. The process according to claim 18, wherein the halogenated promoter is selected from substances capable of generating halogen ions in the reaction solvent.
- of generating bromide ions such as hydrobromic acid and its salts soluble in the reaction medium such as alkaline bromides, ammonium bromide or sodium bromate.
- 15 52. The process according to claim 51, wherein the compound is hydrobromic acid, sodium bromide or potassium bromide.
 - 53. The process according to claim 18, wherein the concentration of halogenated promoter ranges from 0.1 to 50 mg per kg of reaction solvent.
 - 54. The process according to claim 53, wherein the concentration of halogenated promoter ranges from 1 to 10 mg per kg of reaction solvent.
- 55. The process according to claim 18, wherein the reaction is carried out at a temperature ranging from -5

to 90°C.

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56. The process according to claim 55, wherein the temperature ranges from 2 to 50°C.

- 57. The process according to claim 56, wherein the temperature ranges from 20 to 40°C.
- 58. The process according to claim 18, wherein the reaction is carried out at a total pressure higher than atmospheric pressure.
- 59. The process according to claim 58, wherein the total pressure ranges from 10 to 300 bars.
 - 60. The process according to claim 59, wherein the total pressure ranges from 30 to 100 bars.
 - 61. The process according to claim 18, wherein the molar ratio hydrogen/oxygen in the feeding ranges from 1/1 to 1/100.
 - 62. The process according to claim 61, wherein the molar ratio hydrogen/oxygen in the feeding ranges from 1/2 to 1/15.
- 63. The process according to claim 18, wherein the reac-20 tion is carried out in the presence of an inert gas selected from nitrogen, helium, argon.
 - 64. The process according to claim 63, wherein the inert gas is nitrogen.
- 65. The process according to claim 18, wherein the concentration of hydrogen in the gaseous phase in contact

with the reaction solvent is maintained at a value lower than 4.5% molar.

- 66. The process according to claim 18, wherein the reaction is carried out using air as oxygen source.
- 5 67. The process according to claim 18, wherein the reaction is carried out batchwise or in continuous.
 - 68. The process according to claim 18, wherein the solution of hydrogen peroxide is used directly in an oxidation process of a substrate selected from olefins, aromatic hydrocarbons, ammonia and carbonyl compounds, using titanium silicalite as catalyst.
- 69. The process according to claim 18, wherein the solution of hydrogen peroxide is used to obtain aqueous solutions having a commercial titer of H₂O₂ by the removal, from the reaction medium, of the organic components which can be recycled to the synthesis.

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(19) World Intellectual Property Organization
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- 1884 - 1885 | 1885 | 1885 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 1886 | 188

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(54) Title: CATALYST AND ITS USE IN THE SYNTHESIS OF HYDROGEN PEROXIDE

(57) Abstract: The present invention relates to a catalyst consisting of: (a) one or more metals of the platinum group as active components; (b) one or more polyolefins; and (c) a carrier. The invention also relates to a process for the synthesis of hydrogen peroxide (H_2O_2) from hydrogen and oxygen which uses said catalyst and the use of the hydrogen peroxide solution in oxidation processes catalyzed by titanium silicalite. The process operates under high safety conditions with a high productivity and molar selectivity towards the formation of H_2O_2 .

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A. CLASSIFI IPC 7	CATION OF SUBJECT MATTER C01B15/029 C01B21/26 B01J23/42 C07D301/12 C07C249/04 C07D303/04		B01J31/06
According to	international Patent Classification (IPC) or to both national classification	and IPC	
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IPC 7	umentation searched (classification system followed by classification s CO1B BO1J CO7D CO7C		
Documentali	on searched other than minimum documentation to the extent that such	documents are included in	the fields searched
Electronic da	ata base consulted during the international search (name of data base a	THERE COMPEN	UDEY
EPO-Int	ternal, WPI Data, PAJ, CHEM ABS Data,	TNSTEC, COM E	NOL A
C DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Chation of document, with indication, where appropriate, of the relevant	ant passages	Relevant to claim No
X	WO 99 02264 A (MURCHISON CRAIG B; DAVID A (US); WEBB STEVEN P (US); 21 January 1999 (1999-01-21) page 2, line 35 -page 4, line 24 claims 1,8-10,12,14,18	HAYES	1,2,7, 12,18, 45-47, 50-52, 55, 58-60, 66,67 4-6, 8-11, 13-15, 19-25, 48,49, 53,54, 56,57, 61-64
 	and the state of the C	X Patent family mer	nbers are listed in annex.
اشا	orther documents are listed in the continuation of box C.	<u></u>	
"A" docu con. "E" earlik filth "L" docu whl cita "O" docu oth	ment defining the general state of the an which is not sidered to be of particular relevance or document but published on or after the International g date ment which may throw doubts on priority claim(s) or ch is clied to establish the publication date of another tilon or other special reason (as specified) unent referring to an oral disclosure, use, exhibition or er means	or priority date and no cited to understand the invention "X" document of particular cannot be considered involve an inventive s "Y" document of particular cannot be considered cannot be considered.	ed after the International filing date the conflict with the application but the principle or theory underlying the Invertigation of the considered to the power of cannot be considered to the power of cannot be considered to the power of the considered to the considered the considered to the considered the considered the considered the considered the considered the conflict the considered the conflict the
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112		Authorized officer	
Name a	nd malling address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2286 HV Rijswijk Tel. (+31 -70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Rhodes,	K .

Form PCT/ISA/210 (second sheet) (July 1992)

International Application No PCT/EP 02/08546

		PCT/EP 02/08540		
C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	In June 14 and 15 And		
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
Х	US 6 210 651 B1 (HERRMANN WOLFGANG ET AL) 3 April 2001 (2001-04-03) column 1, line 67 -column 4, line 65 claims 1,12-15	1-3,7, 12,16		
Y		4-6, 8-11, 13-15, 19-25, 48,49, 53,54, 56,57, 61-64		
Υ .	WO 98 16463 A (BASF AG ;FLICK KLEMENS (DE); KAIBEL GERD (DE); STAMMER ACHIM (DE);) 23 April 1998 (1998-04-23)	19-25, 48,49, 53,54, 56,57, 61-64		
	claims 1,5-9,12-15 page 4, line 4-16 page 5, line 10-19 page 6, line 26-40 page 8, line 38 -page 9, line 31 examples			
Χ .	EP 0 100 119 A (ANIC SPA) 8 February 1984 (1984-02-08) cited in the application claims	68		
X	US 3 714 342 A (KABISCH G) 30 January 1973 (1973-01-30) claims 1-6	69		
P,A	EP 1 160 196 A (ENI SPA ;ENICHEM SPA (IT)) 5 December 2001 (2001-12-05) the whole document	1-69		
A	US 4 832 938 A (GOSSER LAWRENCE W ET AL) 23 May 1989 (1989-05-23) claims 1-4 column 1, line 48 -column 2, line 2 examples	1-67		
A	EP 0 366 419 A (ATOMIC ENERGY OF CANADA LTD) 2 May 1990 (1990-05-02) the whole document	1-67		
A	EP 0 492 064 A (MITSUBISHI GAS CHEMICAL CO) 1 July 1992 (1992-07-01) column 3, line 3 -column 4, line 4 column 5, line 1-15 examples claims 1-7,10	1-67		

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

nformation on patent family members

Internal	Application No	
PCT/EP	02/08546	

	_				12/ 00348
Patent document cited in search report		Publication date	•	Patent family member(s)	Publication date
WO 9902264	. A	21-01-1999	AU WO	8067598 A 9902264 A1	08-02-1999 21-01-1999
US 6210651	B1	03-04-2001	AU CA CA CN CN EP JP JP WO WO	1994399 A 1994499 A 2315618 A1 2315719 A1 1283165 T 1283166 T 1042220 A1 1042221 A1 2001526167 T 2001526168 T 9932398 A1 9932399 A1 6299852 B1	12-07-1999 12-07-1999 01-07-1999 01-07-1999 07-02-2001 07-02-2001 11-10-2000 11-10-2000 18-12-2001 18-12-2001 01-07-1999 01-07-1999 09-10-2001
WO 9816463	A	23-04-1998	DE AT AU BR DE WO EP JP KR US	19642770 A1 230705 T 725799 B2 5119198 A 9711924 A 59709121 D1 9816463 A1 0946409 A1 2001501904 T 2000049107 A 2001003578 A1	23-04-1998 15-01-2003 19-10-2000 11-05-1998 25-04-2000 13-02-2003 23-04-1998 06-10-1999 13-02-2001 25-07-2000 14-06-2001
EP 0100119	A	08-02-1984	IT AT CA DE DK EP JP JP NO US	1152299 B 21896 T 1196925 A1 3365871 D1 341483 A 0100119 A1 1721832 C 4005028 B 59051273 A 832722 A ,B	31-12-1986 15-09-1986 19-11-1985 09-10-1986 29-01-1984 08-02-1984 24-12-1992 30-01-1992 24-03-1984 3, 30-01-1984 23-05-1989
US 3714342	· A	30-01-197	DE AT AT BE CA CH CS ES FR BNL PL SE TR	219470 A1	19-05-1971 10-05-1978 15-09-1977 15-02-1971 09-10-1973 15-01-1975 23-08-1974 01-01-1973 25-06-1971 08-08-1973 12-03-1971 30-08-1975 27-01-1975 02-05-1976 28-02-1982 28-04-1971

Form PCT/ISA/210 (patent family annex) (July 1992)

information on patent family members

International Application No PCT/EP 02/08546

•				
Patent document cited in search report .	Publication date		Patent family member(s)	Publication date
EP 1160196 A	. 05-12-2001	IT	MI20001219 A1	03-12-2001
El 1100130 A	. 05 12 2001	ĒΡ	1160196 A1	05-12-2001
		ĴΡ	2002029711 A	29-01-2002
		UŞ	2002028174 A1	07-03-2002
US 4832938 A	23-05-1989	AU	608976 B2	18-04-1991
03 4032300	20 00 2005	ΑÜ	3476089 A	16-11-1989
	•	BR	8902234 A	09-01-1990
•		CA	1312720 A1	19-01-1993
	•	CN	1037874 A ,B	13-12-1989
•	•	EP	0342048 A2	15-11-1989
		ΝZ	229087 A	28-08-1990
•		PT	90542. A	30-11-1989
		TR	24229 A	03-07-1991
EP 0366419 A	02-05-1990	CA	1317740 A1	18-05-1993
EP 0366419 A	02-05-1990	DE	68917255 D1	08-09-1994
		DE	.68917255 T2	19-01-1995
		. EP	0366419 A1	02-05-1990
		JP	. 2258610 A	19-10-1990
		US	5082647 A	21-01-1992
		03	3002047 A	21-01-1992
EP 0492064 A	01-07-1992	CA	2052009 A1	28-06-1992
Li 0452004 /		DE	69101775 D1	26-05-1994
•	•	DE	69101775 T2	22-09-1994
•	•	DE	492064 T1	05-11-1992
		EP	0492064 A1	01-07-1992
		JΡ	4357105 A	10-12-1992
		ÜS	5132099 A	21-07-1992

THIS PAGE BLANK (USPTO)